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| | |
|----------------|--|
| NEWS 1 | Web Page for STN Seminar Schedule - N. America |
| NEWS 2 DEC 01 | ChemPort single article sales feature unavailable |
| NEWS 3 JAN 06 | The retention policy for unread STNmail messages will change in 2009 for STN-Columbus and STN-Tokyo |
| NEWS 4 JAN 07 | WPIDS, WINDEX, and WPIX enhanced Japanese Patent Classification Data |
| NEWS 5 FEB 02 | Simultaneous left and right truncation (SLART) added for CERAB, COMPUAB, ELCOM, and SOLIDSTATE |
| NEWS 6 FEB 02 | GENBANK enhanced with SET PLURALS and SET SPELLING |
| NEWS 7 FEB 06 | Patent sequence location (PSL) data added to USGENE |
| NEWS 8 FEB 10 | COMPENDEX reloaded and enhanced |
| NEWS 9 FEB 11 | WTEXTILES reloaded and enhanced |
| NEWS 10 FEB 19 | New patent-examiner citations in 300,000 CA/CAPplus patent records provide insights into related prior art |
| NEWS 11 FEB 19 | Increase the precision of your patent queries -- use terms from the IPC Thesaurus, Version 2009.01 |
| NEWS 12 FEB 23 | Several formats for image display and print options discontinued in USPATFULL and USPAT2 |
| NEWS 13 FEB 23 | MEDLINE now offers more precise author group fields and 2009 MeSH terms |
| NEWS 14 FEB 23 | TOXCENTRE updates mirror those of MEDLINE - more precise author group fields and 2009 MeSH terms |
| NEWS 15 FEB 23 | Three million new patent records blast AEROSPACE into STN patent clusters |
| NEWS 16 FEB 25 | USGENE enhanced with patent family and legal status display data from INPADOCDB |
| NEWS 17 MAR 06 | INPADOCDB and INPAFAMDB enhanced with new display formats |
| NEWS 18 MAR 11 | EPFULL backfile enhanced with additional full-text applications and grants |
| NEWS 19 MAR 11 | ESBIOBASE reloaded and enhanced |
| NEWS 20 MAR 20 | CAS databases on STN enhanced with new super role for nanomaterial substances |
| NEWS 21 MAR 23 | CA/CAPplus enhanced with more than 250,000 patent equivalents from China |
| NEWS 22 MAR 30 | IMSPATENTS reloaded and enhanced |
| NEWS 23 APR 03 | CAS coverage of exemplified prophetic substances enhanced |
| NEWS 24 APR 07 | STN is raising the limits on saved answers |

NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3,
AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

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NEWS LOGIN Welcome Banner and News Items

NEWS IPC8 For general information regarding STN implementation of IPC 8

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* *

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=> fil reg
COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 0.22 0.22

FILE 'REGISTRY' ENTERED AT 08:31:15 ON 15 APR 2009
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STRUCTURE FILE UPDATES: 14 APR 2009 HIGHEST RN 1134418-75-9
DICTIONARY FILE UPDATES: 14 APR 2009 HIGHEST RN 1134418-75-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

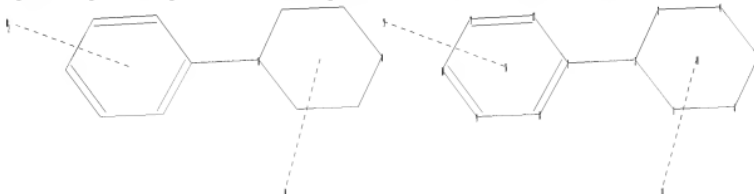
TSCA INFORMATION NOW CURRENT THROUGH January 9, 2009.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

<http://www.cas.org/support/stngen/stndoc/properties.html>

=>
Uploading C:\Program Files\Stnexp\Queries\QUERIES\105678481.str



chain nodes :

```

13 15
ring nodes :
1 2 3 4 5 6 7 8 9 10 11 12
chain bonds :
2-7
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-12 8-9 9-10 10-11 11-12
exact/norm bonds :
1-2 1-6 2-3 2-7 3-4 4-5 5-6
normalized bonds :
7-8 7-12 8-9 9-10 10-11 11-12

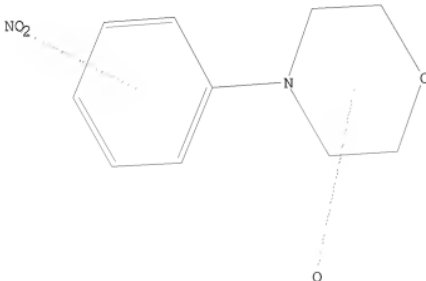
Match level :
1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS

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L1 STRUCTURE UPLOADED

→ d

L1 HAS NO ANSWERS



Structure attributes must be viewed using STN Express query preparation.

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=> s 11  
SAMPLE SEARCH INITIATED  
SAMPLE SCREEN SEARCH
```

3 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
PROJECTED ANSWERS: BATCH **COMPLETE**
PROJECTED ITERATIONS: 15954 TO 19526
PROJECTED ANSWERS: 3 TO 1562

13 3 SEA SCS SAM 11

=> s.11.full

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FULL SCREEN SEARCH COMPLETED - 17886 TO ITERATE

100.0% PROCESSED 17886 ITERATIONS
SEARCH TIME: 00.00.01

50 ANSWERS

L3 50 SEA SSS FUL L1

=> s l3 and caplus/lc
65173951 CAPLUS/LC
L4 36 L3 AND CAPLUS/LC

=> s l3 not l4
L5 14 L3 NOT L4

=> d 15 1-14

L5 ANMERK 1 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN

STN 899135-53-3 REGISTRY

ED Entered STN: 07 Aug 2006

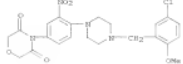
CH INDEX NAME NOT YET ASSIGNED

MF C22 H21 Cl N4 O6

SR Chemical Library

Supplier: Aurora Fine Chemicals

LC STN File#: C8EMCAG75



PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

L5 ANMERK 2 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN

STN 899135-43-8 REGISTRY

ED Entered STN: 07 Aug 2006

CH 3,5-Morpholinolindole, 4-[4-(4-methoxyphenyl)-1-piperazinyl]-3-

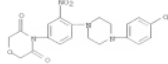
nitrophenyl]- (CA INDEX NAME)

MF C22 H21 Cl N4 O6

SR Chemical Library

Supplier: Aurora Fine Chemicals

LC STN File#: C8EMCAG75



PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

L5 ANMERK 3 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN

STN 899135-44-9 REGISTRY

ED Entered STN: 07 Aug 2006

CH 3,5-Morpholinolindole, 4-[4-(3,6-dihydro-4-phenyl-1(2H)-pyridinyl)-3-

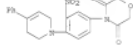
nitrophenyl]- (CA INDEX NAME)

MF C22 H21 N4 O6

SR Chemical Library

Supplier: Aurora Fine Chemicals

LC STN File#: C8EMCAG75



PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

L5 ANMERK 4 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN

STN 899135-45-0 REGISTRY

ED Entered STN: 07 Aug 2006

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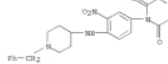
piperazinylmethyl)phenyl]- (CA INDEX NAME)

MF C22 H21 N4 O5

SR Chemical Library

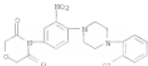
Supplier: Aurora Fine Chemicals

LC STN File#: C8EMCAG75



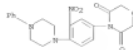
PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

L5 ANMERK 5 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 693735-20-1 REGISTRY
 ED Entered STN: 22 Jun 2004
 CH 3,5-Morpholinodione, 4-[4-(2-chlorophenyl)-1-piperazinyl]-3-nitrophenyl]-
 (CA INDEX NAME)
 MF C22 H22 N2 O5
 SR Chemical Library
 Supplier: Aurora Fine Chemicals
 LC STN File#: CEMCATS



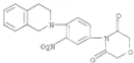
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 RS 693735-40-6 REGISTRY
 ED Entered STN: 22 Jun 2004
 CH 3,5-Morpholinodione, 4-[4-(4-phenyl-1-piperazinyl)phenyl]-
 (CA INDEX NAME)
 MF C22 H22 N2 O5
 SR Chemical Library
 Supplier: Interchim
 LC STN File#: CEMCATS



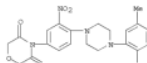
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 7 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 693735-41-0 REGISTRY
 ED Entered STN: 22 Jun 2004
 CH 3,5-Morpholinodione,
 4-[4-(2,5-dimethylphenyl)-1-isoquinolinalyl]-3-nitrophenyl]-
 (CA INDEX NAME)
 MF C22 H24 N2 O5
 SR Chemical Library
 Supplier: ChemDiv, Inc.



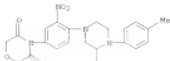
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 8 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 693735-42-4 REGISTRY
 ED Entered STN: 22 Jun 2004
 CH 3,5-Morpholinodione, 4-[4-(4-(2,5-dimethylphenyl)-1-piperazinyl)-3-
 nitrophenyl]-
 (CA INDEX NAME)
 MF C22 H24 N4 O5
 SR Chemical Library
 Supplier: ChemDiv, Inc.
 LC STN File#: CEMCATS



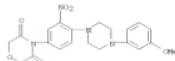
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 9 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 697293-93-9 REGISTRY
 ED 2004-06-14 Entered 2004
 CH 3,5-Morpholinedione, 4-[4-(3-methyl-4-(4-methylphenyl)-1-piperazinyl)-3-nitrophenyl]- (CA INDEX NAME)
 MF C21 H27 N5 O6
 SR Chemical Library
 Supplier: Chem3Dv, Inc.
 LC STN File#: CHEMCATS



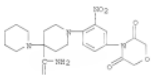
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 10 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 697293-92-8 REGISTRY
 ED 2004-06-14 Entered 2004
 CH 3,5-Morpholinedione, 4-[4-(3-methoxyphenyl)-1-piperazinyl]-3-nitrophenyl]- (CA INDEX NAME)
 MF C21 H27 N5 O6
 SR Chemical Library
 Supplier: Chem3Dv, Inc.
 LC STN File#: CHEMCATS



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 11 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 697293-94-0 REGISTRY
 ED 2004-06-14 Entered 2004
 CH [1,4'-Bis[perimidin]-4'-imino]anabacone,
 [1,4'-Bis[perimidin]-4'-imino]anabacone-4-nitrophenyl]- (CA INDEX NAME)
 MF C21 H27 N5 O6
 SR Chemical Library
 Supplier: Chem3Dv, Inc.
 LC STN File#: CHEMCATS



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANMERK 12 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
 RS 697293-95-1 REGISTRY
 ED 2004-06-14 Entered 2004
 CH 3,5-Morpholinedione, 4-(3-nitrophenyl)- (CA INDEX NAME)
 MF C16 H14 N2 O4
 SR Chemical Library
 Supplier: Scientific Exchange, Inc.
 LC STN File#: CHEMCATS



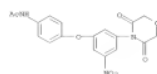
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

L5 ANNUAL 13 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
RN 341619-43-0 REGISTRY
RD 2009-01-01 2009
CN 2,5-Morpholinedione, 4-(4-fluoro-3-nitrophenyl)- ICA INDEX NAME
MW C14 H11 F NO2
SB Chemical Library
Supplier: Scientific Exchange, Inc.
LC STN Fileas: C8H6NO2



PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

L5 ANNUAL 14 OF 14 REGISTRY COPYRIGHT 2009 ACS on STN
RN 313249-13-7 REGISTRY
RD 2009-01-01 2009
CN Acetanamide, N-[4-[(3,5-dioxo-4-morpholinyl)-5-nitrophenyl]phenyl]- ICA
INDEX NAME
MW C18 H14 N2 O3
SB Chemical Library
Supplier: Nanosys Combinatorial Synthesis Inc.
LC STN Fileas: C8H14NO2



PROPERTY DATA AVAILABLE IN THE "PROP" FORMAT

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COST IN U.S. DOLLARS          SINCE FILE      TOTAL
                                ENTRY        SESSION
FULL ESTIMATED COST          220.41       220.63
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FILE COVERS 1907 - 15 Apr 2009 VOL 150 ISS 16
FILE LAST UPDATED: 14 Apr 2009 (20090414/ED)

Caplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

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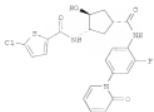
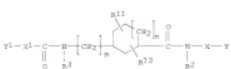
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L1      3 S L1
L2      50 S L1 FULL
L3      36 S L3 AND CAPLUS/LC
L4      14 S L3 NOT L4
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FILE 'CAPLUS' ENTERED AT 08:32:28 ON 15 APR 2009

=> s 14

L6 36 L4

=> d ibib abs hitstr 1-36



AB Title compds. I [X1 = carboxyl, cyano, alkoxycarbonyl, etc.; X2 = H or a member selected from -CH2-, -cycloalkyl, heterocyclyl, etc. (wherein one or two carbon atoms of the heterocyclyl are optionally substituted by one or more substituents selected from R4; X3 = arylene, heteroarylene or heterocyclylene (wherein the arylene, heteroarylene and heterocyclylene are optionally substituted by one or more substituents selected from R4); R3 = alkyl, heteroalkyl or heterocyclyl (where the alkyl, heteroalkyl and heterocyclyl are optionally substituted by one or more substituents selected from halogen, cyano, nitro, etc.); Y1 = H, aryl, heteroaryl, etc. (wherein the aryl and heteroaryl are substituted by one or more substituents selected from halogen, cyano, nitro, etc.)]; n = 0, 1 or 2]; products and pharmaceutically useful compositions comprising the same. Preparation of [12,25,43]-N-(300'-1-amino-2-hydroxyazepatane-4-carboxylic acid Me ester followed by EGP mediated cyclization of 5-chloro-2-thiophenecarboxylic acid (5-chlorothiophene-2-carboxylic acid) in the presence of trimethylaluminum and compound II. In coagulation factor Xa (FXa) inhibition assays, the K1 value of compound II was 0.015 μM.

IT K17870-44-0 (Synthetic preparation); PAPF (Preparation); RACT (Reaction or reagent); EGM (Synthetic preparation); containing thiophene compds. containing cyclopentanecarboxamide moiety

K17870-44-0 CAPLUS

CG Benzodiazepine, 2-oxo-5-(3-oxo-4-morpholinyl)- (CA INDEX NAME)



ACCESSION NUMBER: 1461274374

DOCUMENT NUMBER: 1461274374

TITLE: Preparation of five-membered aromatic heterocycles,

their salts and their pharmaceutical use

INVENTOR(S): Taniguchi, Takahiko; Fujimoto, Takeshi; Tokushita,

Hidemaru; Tashiro, Shigeo

THE NIPPON DRUG COMPANY LTD., Ltd., Japan

PATENT ASSIGNEE(S): Jpn.: Kokai Tokkyo Koho, 64pp.

SOURCE: CA/CN/JPO/JAIS

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

PARENT DOCUMENT COUNT: 1

PATENT INFORMATION:

PATENT NO.: 20070245752

KIND: A

DATE: 20070222

APPLICATION NO.: JP 2005-232392

DATE: 20050810

PROVISIONAL APPLICATION INFO.: JP 2005-232392

DATE: 20050810

OTHER SOURCE(S): MOJCAT 1461274374

GS



AB Title compds. I [X1 = bond, (un)substituted Cl-4 alkenyl; Y = NHCO, NHCOOMe, NHCOMe, Zn, Zn-alkyl, Zn-(un)substituted Cl-4 alkyln, n = 0-2; Z, Z' = bond, (un)substituted Cl-4 alkenyl, (un)substituted Cl-4 alkenylene, (un)substituted Cl-4 alkenylene, etc.; X2 = (un)substituted aryl, (un)substituted heterocyclic; D, E = (un)substituted cyclic group, ring A = 5-membered aromatic heterocycle] or their salts are prepared. The compounds are useful as long-lasting oral anticoagulants for treatment of myocardial infarction, stroke, thromboembolic disease, etc. Thus, gefitinib, 5-chloro-2-(3-oxo-1-phenylpropyl)-2-methoxy-2-nitrobenzoate with 4-(4-azidomethyl)morpholin-3-one in MeOH gave 34% yield of 4-[4-(4-azidomethyl)morpholin-3-one-3-oxo-1-phenylpropyl]-2-methoxy-2-nitrobenzoate, which inhibited human FXa with IC50 value of 2.1 nM.

IT K178729-44-0 (Synthetic preparation); PAPF (Preparation); RACT (Reaction or reagent); EGM (Synthetic preparation); containing preparation of aromatic heterocycles as blood coagulation factor Xa inhibitors)

K178729-44-0 CAPLUS

CG 4-(3-methyl-4-nitrophenyl)-, 4-(3-methyl-4-nitrophenyl)- (CA INDEX NAME)



ACCESSION NUMBER: 2005159650 CARLOS

DOCUMENT NUMBER: 142-241542 CARLOS

TITLE: Method for production of *M*-arylmorpholines from 5-chloro-2,3-dihydro-1,4-dioxin

INVENTOR(S): Dabholkar, Suresh; Mahrer, Werner; Medek, Hans

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd.; Werner, Mahrer & Partner G.m.b.H., Germany

SOURCE: PCT Int. Appl. 04 49 pp.

DOCUMENT TYPE: PCT Int. Appl.

FAMILY ACC. NUM. COUNT: 1 CARLOS

PATENT INFORMATION:

PATENT NO.: WO 2005014899 A1

CITE NO.: 2005014899A1

DATE: 200501224

APPLICATION NO.: WO 2004-EP7938

DATE: 2004-09-23

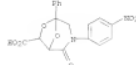
W0 2005014899 A1

WO 2005014899 A1

Reaction of 3-oxa-6,8-dioxabicyclo[3.2.1]octane-7-carboxylates and analogs

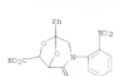
RN: 531467-21-3 CAPLUS

CH: 6,8-Dioxa-3-azabicyclo[3.2.1]octane-7-carboxylic acid,
3-(4-nitrophenyl)-2-oxo-5-phenyl- (CA INDEX NAME)



RN: 351467-28-8 CAPLUS

CH: 6,8-Dioxa-3-azabicyclo[3.2.1]octane-7-carboxylic acid,
3-(4-nitrophenyl)-2-oxo-5-phenyl- (CA INDEX NAME)



REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

TOMCAT

TOMCAT

andled, Gennar Martin, M. Victoria; Parada, N.; Garcia, Instituto de Química Orgánica General, C.S.I.C.-

Biotecnología, Madrid, Spain; Instituto de Heterociclos (2000), 52(1), 237-251

Coden: KFUTAM; ISSN: 0255-5414

Journal: Journal of the Institute of Heterocyclic Chemistry

Language: ENGLISH

Other Source: CHEMABS 112:194237

AB: The behavior of the 2(5H)-furanones towards p-methoxy- and p-nitrophenyl azides has been investigated, in particular with respect to the regio-

and stereoselectivity. The 1,3-dipolar cycloaddn. of the azomethine ylide generated from 2(5H)-furanone and trans-1-(p-methoxyphenyl)aziridine-2,3-dianhydride to 2(5H)-furanones proceeds in good yield and affords functionalized furol[3,4-*p*]-cyclopent-1-one derivatives.

17 DOCUMENT NUMBER: 255728-49-7 CAPLUS

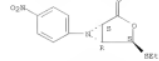
EL: IUPAC (Synthetic preparation); PEER (Preparation)

Reaction of 2(5H)-furanones, of aryl azides or an aziridine ylide with furanones

RN: 255728-49-7 CAPLUS

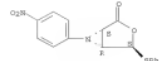
CH: 3-Oxo-4-azabicyclo[1.1.0]hexan-2-one, 4-(ethylthio)-6-(4-nitrophenyl)-, (1R,4S,5S)-rel- (CA INDEX NAME)

Relative stereochemistry:



RN: 255728-49-7 CAPLUS
CH: 3-Oxo-4-azabicyclo[1.1.0]hexan-2-one, 6-(4-nitrophenyl)-4-(phenylthio)-, (1R,4S,5S)-rel- (CA INDEX NAME)

Relative stereochemistry:



REFERENCE COUNT:

30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR

TRIS

AUTHOR(S): Kostyuk, V. V.; Levin, A. A.; Shein, Yu. D.; Ionin, I.

B. I.; Stilkina, E. F.; Karchevskaya, L. A.; Chukunova, E. V.; Matrenikova, R. S.; Shneider, M. A.; Vass. Nauchno-Issled. Tekhnol. Inst. Antibiol. i Virolog. im. N. S. Klyushnikova, Moscow, USSR

SOURCE: Khimiko-Farmatsevticheskii Zhurnal (1990), 24(9), Khimiko-Farmatsevticheskii Zhurnal (1990), 24(9), 114-120; CHEMABS 114:02381

DOCUMENT TYPE: Journal

LANGUAGE: Russian

OTHER SOURCE(S): CHEMTRACT 114:02381

GI



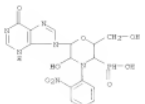
AB: The title virucidal phosphonate derivative I [D = carboxybutyl, α -CH2CH2CH2CH3, β -CH2CH2CH2CH3] was synthesized and its antiviral properties were studied. Compound I was found to be active against HSV-1 and HSV-2 and was also found to be active against VZV and varicella-zoster virus.

17 DOCUMENT NUMBER: 255728-50-8 CAPLUS
EL: IUPAC (Biological activity or effector, except adverse); RU (Biological activity or effector, except adverse); SRW (Synthetic preparation); SRW (Biological activity or effector, except adverse); PEER (Preparation)

Reaction of 2(5H)-furanones with 2(5H)-furanones

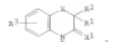
RN: 112019-63-3 CAPLUS

CH: Phosphonic acid, 6-[1,4-dihydro-6-oxo-2-pyran-3-yl]-5-hydroxy-2-(hydroxybutyl)-4-(4-nitrophenyl)-2-naphthalenyl- (CA INDEX NAME)



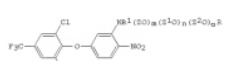
L6 ANSWER 34 OF 36 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 198718463 CAPLUS
DOCUMENT NUMBER: 106-18463
ORIGINAL REFERENCE NO.: 106-13165a, 3172a
TITLE: Synthesis and anthelmintic activity of some new 6

and
 7-azabicyclo[2.1.0]hex-1-yl-benzo[1,2-d]thiophene-2-carboxylic acid (7a)
 and benzoxazolo[3,2-d]thiophene-2-carboxylic acid (7b).
AUTHOR(S): K. P. Rao, K. Srinivasan Singh, A. N. Jagtap,
 K. V. Rao, M. J. Jain, L. Gaudelli, S. G. Krishnamoorthy, S. R. Rao, S. S. Venkateswaran, S. S. Rao,
CORPORATE SOURCE: IITM Research Foundation, IITM,
 Hesarappa, IITM, Hyderabad, 500 037, India
SOURCE: Indian Journal of Chemistry, Section B: Organic
 Chemistry including Medicinal Chemistry (1995),
 34(12), 1121-1126.
CODEN: IJCBAB
DOCUMENT TYPE: Journal
LANGUAGE: English
NUMBER OF PAGES: 12
PUBLISHER: IITM, Hyderabad
ISSN: 0377-0428
DOI: 10.1002/jocb.5460341203

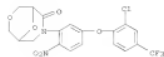


16. ASHEM 35 OF 34 CAJUS COUPRAGHT 2009 ACS ON STM
1986-144844 CAJUS
DOCUMENT NUMBER: 104-148484
ORIGINAL REFERENCE NO.: 104-148484, 24488a
TITLE: Polyalkylbenzimidophenyl ethers having pesticidal activity
INVENTOR(S): Danner, Dieter
PATENT ASSIGNEE(S): Ciba-Geigy AG, Switzerland
SOURCE: Eur. Pat. Appl., 38 pp.
COUNTRIES: EP/DMX
PUBLICATION DATE: 1986-02-05
LANGUAGE: German
FAMILY ACC. NEW COUNT: 1

| PATENT INFORMATION: | | | |
|-----------------------------------|------|----------|-----------------|
| PATENT NO. | KIND | DATE | APPLICATION NO. |
| EP 149427 | A1 | 19850724 | EP 1984-810620 |
| EP 149427 | A | 19870708 | |
| R: BE, CH, DE, FR, GB, IT, LI, NL | | | |
| US 4494105 | A | 19870915 | US 1984-679860 |
| JP 60172549 | A | 19850906 | JP 1984-249595 |



AS The title compound, (I_2 : $R =$ alkyl, alkylidene, etc.; $E_1 =$ H , alkyl, hydroxylalkyl, alkoxyalkyl, etc.; $E_2 =$ H , Cl, F, Z_1 , $Z_2 =$ X -alkyl; $n_1, n_2 = 0-1$) was prepared as herbicides, insecticides, and acaracides, and typically prepared by condensation of the corresponding substituted phenylhydrazine derivative with a substituted benzoyl derivative with the pertinent polyisoprene in the presence of an anti-oxidation agent. Thus, the condensation of I_2 with $\text{HOC(=O)CH}_2\text{COCH}_3$ in Me_2CO , over $2-\text{butoxyphenylhydrazine}-2'-\text{chloro}-4-\text{nitro}-4'-\text{trifluoromethylphenyl ether}$ (II), gave the title compound, which was found to be active against controlled species, *Solanum tuberosum*, *Spiraea alba* and *Stellaria media*.
 10129-68-19
 (10129-68-19)
 (herbicide preparation); PFR (Preparation)
 (preparation of, as herbicide and insecticide)
 10129-68-19-CARBO
 (10129-68-19-CARBO)
 7-[5-[2-chloro-4-(2-[trifluoromethyl]phenoxyl)-2-nitrophenyl]-3-(Z -CA INDEX



16 AMERICA 35 OF 36 CAPLUS COPYRIGHT 2009 ACS on STN

1972-11-21 70 CAPLUS

DOCUMENT NUMBER:

DOI NUMBER:

ORIGINAL PREFERENCE NO.:

TITLE:

Conjugated systems obtained by reaction of cyclic
anhydrides with nucleophilic reagents and dehydrating agents.
III. Macrocyclic compounds. Anhydride dihydroxides of
1,4-dialkyl-1,2,3,5-tetra(aryloxy)-2,4-

1,4-dihydropyrazines.

Bott, M.; Bouček, J.

Inst. Macromol. Chem., Czech. Acad. Sci., Prague,
Czechoslovakia

SOURCE:

Tetrahedron (1972), 28(15), 607-10

ISSN: 0040-4020

DOCUMENT TYPE:

Journal

LANGUAGE:

English

DOI NUMBER:

10.1016/S0040-4020(00)89925-5

01 For diagram(s), see printed CA issue.

AS Alternative name: 1,4-Bis(phenylmethoxy)-2,4-dihydropyrazine.

AS Alternative name: 1,4-Bis(phenylmethoxy)-2,4-dihydropyrazine with H atoms at the para positions of the Ph rings systematically substituted with a NO₂ group, Br and a NO group

substitutive; of the same compound with Ph groups systematically

substituted with NO₂ groups at positions 1 and 4 were prepared. The 1,4-NMe₂ and1,4-Br₂ derivatives of these compounds are in agreement with the assumed prevalence participation of an aromatic enone structure (II) in their real structure.

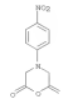
17 26(15), 607-10

EIC: EMM (Synthetic preparation); PREP (Preparation)

(printed page 607)

356-34-5 CAPLUS

CH 2,4-Morpholinodione, 4-(4-nitrophenyl)- (CA INDEX NAME)



=> log y

| | | |
|--|------------|---------|
| COST IN U.S. DOLLARS | SINCE FILE | TOTAL |
| FULL ESTIMATED COST | ENTRY | SESSION |
| | 204.54 | 425.17 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE | TOTAL |
| CA SUBSCRIBER PRICE | ENTRY | SESSION |
| | -29.52 | -29.52 |

STN INTERNATIONAL LOGOFF AT 08:34:00 ON 15 APR 2009